Effect of Pit and Fissure Sealants on the Prevention of Enamel Demineralization After Exposure to Streptococcus mutans Biofilm: An In Vitro Study

Patrícia da Silva Lopes Pereira da Silva¹ Maristela Barbosa Portela¹ Antônio Ferreira-Pereira²
Mônica Almeida Tostes¹

¹Department of Pediatric Dentistry, School of Dentistry, Universidade Federal Fluminense, Niterói, RJ, Brazil
²General Microbiology Department, Paulo de Góes Microbiology Institute, Universidade Federal do Rio de Janeiro, Rio de Janeiro, RJ, Brazil

How to cite this article: Pereira da Silva PSL, Portela MB, Ferreira-Pereira A, Tostes MA. Effect of Pit and Fissure Sealants on the Prevention of Enamel Demineralization After Exposure to Streptococcus mutans Biofilm: An In Vitro Study. J Pediatr Dent 2020;6(1):12-19

Abstract

Objective: The aim of this study was to evaluate the effect of sealants on the prevention of enamel demineralization and on biofilm metabolic activity.

Materials and Methods: Cavity preparations were performed on 45 blocks of bovine teeth (4x4x4 mm) randomly assigned to three groups (n=15): RI-Riva Light Cure®/SDI; EM-Embrace™ WetBond™ Pulpdent Corp.®; and CO-Natural Flow/DFL resin (negative control). The sealed blocks were subjected to thermocycling (500 cycles/30 s). Half of the blocks were covered with acid-resistant varnish to create a control area (RI, EM, and CO) and an experimental area (RI-EX, EM-EX, and CO-EX). They were subsequently exposed to Streptococcus mutans biofilm for assessment of demineralization by the Knoop microhardness test (50 g/15 s), at 25, 50, 100, 150, and 200 µm from the interface. In another test, the cylinders of the sealants (4 mmx3 mm) were subjected to the biofilm metabolic activity test.

Results: The RI-EX group showed higher microhardness than CO-EX (p<0.05) and less demineralization when compared to RI at all distances. Microbial activity was lower in EM compared to CO, but not statistically significant in relation to RI.

Conclusion: Both sealants can inhibit enamel demineralization in the presence of S. mutans biofilm.

Keywords: Dental enamel, dental resin, hardness test, glass ionomer cements

Introduction

Routine dental care is sought out to prevent caries and to arrest its progression.[1,2] Dental caries has a high prevalence in children and it is a public health problem.[3] The use of pit and fissure sealants has been well supported for preventing dental caries or arresting its progression on occlusal surfaces of both permanent and primary teeth.[2,4,5] Fluoride release by sealant materials has been suggested to reduce the adhesion of S. mutans and may therefore contribute to caries prevention.[6–9] Previous studies have reported that fluoride level in Embrace WetBond (EWB) and glass ionomer showed higher fluoride release in water
and artificial saliva and had greater antimicrobial activity than did resin sealant. Clinically, no difference was noted in the anticaries effect between ionomer and EWB sealants. However, despite many comparative studies on resin sealant and ionomer material, only limited information is available on Riva Light Cure® and Embrace sealants applied on fissures. Embrace sealants are biocompatible, do not contain bisphenol A, have hydrophilic properties, in addition to fluoride release, and have been indicated in pediatric dentistry. Notwithstanding the anticaries efficacy of sealant treatment, the results remain inconsistent.

Accordingly, the present study evaluated the effect of an ionomer, Riva light Cure® sealant, and EWB on the prevention of enamel demineralization after *S. mutans* biofilm formation in vitro.

**Materials and Methods**

The samples were subjected to thermocycling, cariogenic biofilm formation, and Knoop microhardness test, and the cylinders of the sealants were subjected to a cariogenic biofilm metabolic activity test. The study design is shown in Fig. 1.

**Preparation of the specimens**

Sample size was calculated by establishing a statistical power of 0.95 and significance of 0.05 with effect size 1. At least 13 specimens were necessary for each group. The bovine teeth were kept in a 10% formaldehyde solution until use. The teeth were polished at low speed with an Abbott-Robinson® bristle brush using pumice mixed with distilled water (Kavo Dental GmbH, Biberack, Germany) and analyzed thereafter under a stereoscopic magnifying glass (CGA 6745, Tecnival, Buenos Aires, Argentina). Only those teeth without cracks or macroscopic defects or any other enamel alterations were selected.

The crowns were separated from the roots with a double-sided diamond disc (Diamond wafering blades ½” diameter 4” x 0.012” 7–ref 11–4244, Buehler Ltda, Lake Bluff, IL, USA). Buccal surfaces were separated using a water-cooled diamond-impregnated low-speed saw. An enamel block was obtained from each tooth. After embedding the blocks in acrylic resin, the buccal surfaces of the enamel specimens (4 mm x 4 mm x 4 mm) were ground with SiC paper (400, 600, and 1.200 grit sizes) to obtain flat surfaces. Each specimen was flattened and polished before cavity preparation. The 45 enamel blocks were therefore fixed with wax at the center of a crystal acrylic base (30 mm in diameter and 8 mm in thickness), with the largest flat area of the enamel facing downwards. A 320-grit sandpaper (Carbimet Paper discs, Buehler, Lake Buff, IL, USA) was used under water cooling for dentin planing. The blocks were fixed with the embedded enamel surfaces so that they were visible, but they were flattened at low speed for 1 min using Politriz (Arotec, Cotia, SP, Brazil) and 600, 1000, and 1200-grit sandpapers (Extec Corp. Enfield, CT, USA) and for 2 min using a 2500-grit sandpaper (Extec Corp. Enfield, CT, USA). The whole sanding process was carried out manually and the samples were etched under minimum pressure to keep them on the sander plate.

**Cavity preparation**

A standardized cavity preparation was performed on the enamel of each block. The blocks were attached to acrylic bases in the central buccal region where a cavity was formed using a 2292 KG Sorensen diamond drill (Barueri, SP, Brazil) at high speed under generous air/water cooling. A “stop” mechanism was employed so that the cavity preparation pattern was maintained for every 10 teeth. After that, prophylaxis with pumice and water paste, application of 37% phosphoric acid for 30 s, and washing with water for 60 s were performed. The restorations were performed according to the manufac-
turer’s instructions (Table 1). The samples were polished with 3000-grit sandpaper (3M, Maplewood, Minnesota, USA) and 1-µm Alumina powder (Buehler, Lake Buff, IL, USA) and finally polished with felt disc (TEXMET C, ref. 40–1108, Buehler Ltda, Lake Buff, IL, USA) for 3 min at high speed until a glassy surface was obtained. At the end, the samples were observed under an optical microscope (Stemi 2000C, Zeiss, Jena, Germany) with a 10X objective until no material covered the tooth margin.

Thermocycling procedure

The samples were kept for 24 h in distilled water and placed in a thermocycler (Marconi, MA 470/3C, Piracicaba, SP, Brazil) at 5°C, 37°C, and 55°C in 500 cycles of 30 s in each bath, considering a transfer time of 10 s between baths. The samples were subjected to S. mutans biofilm activity as described below.

Induction of S. mutans biofilm formation

The blocks were covered with two layers of acid-resistant enamel (Colorama), except for half of the restored buccal surface (Fig. 1). No excess sealant had been observed at the preparation margin (optical microscope, 10x). Thus, half of the blocks, including enamel and sealant, were covered with acid-resistant varnish to create a control area (RI, EM, and CO) and an experimental area (RI-EX, EM-EX, and CO-EX), the latter of which was exposed to the biofilm.

In this phase, the samples were fixed in 45 wells (subdivided into 15 per group), in acrylic plates (TPP 24 ZellkulturTestplatte F, Trasadingen, Switzerland), and sterilized by ethylene oxide. The

---

Table 1. Composition, indication, and insertion technique of the tested materials

<table>
<thead>
<tr>
<th>Dental materials</th>
<th>Composition</th>
<th>Indication</th>
<th>Insertion Technique</th>
</tr>
</thead>
<tbody>
<tr>
<td>Riva light® cure</td>
<td>Resin-reinforced glass ionomer cement.</td>
<td>II/II smaller restorations, primary and geriatric teeth; core constructions; root surface restorations; filling or padding of temporary restorations; sealing of pits and fissures.</td>
<td>Prophylaxis with pumice and water, 37% phosphoric acid etching for 10 s; washing and drying with light air jet; mixing for 10 s in Ultramat S; adaptation for the Riva applicator; photoactivation for 20 s.</td>
</tr>
<tr>
<td>Embrace™ Wetbond™</td>
<td>BisGMA, dimethacrylates, aluminum amorphous silica and 36.6% of charge.</td>
<td>III/IV class restorations; filling of regions; cementation of porcelain veneers; bonding of fragments; sealing of implant screw hole; repair of resin and porcelain erosion.</td>
<td>Prophylaxis with pumice and water; application of resin in a single increment; photoactivation for 30 s.</td>
</tr>
<tr>
<td>Natural flow resin</td>
<td>Natural flow resin.</td>
<td>II/III smaller restorations, primary and geriatric teeth; core constructions; root surface restorations; filling or padding of temporary restorations; sealing of pits and fissures.</td>
<td>Prophylaxis with pumice and water; 37% phosphoric acid etching for 30 s; washing and drying with light air jet; sealing of pits and fissures.</td>
</tr>
</tbody>
</table>

---

The samples were polished with 3000-grit sandpaper (3M, Maplewood, Minnesota, USA) and 1-µm Alumina powder (Buehler, Lake Buff, IL, USA) and finally polished with felt disc (TEXMET C, ref. 40–1108, Buehler Ltda, Lake Buff, IL, USA) for 3 min at high speed until a glassy surface was obtained. At the end, the samples were observed under an optical microscope (Stemi 2000C, Zeiss, Jena, Germany) with a 10X objective until no material covered the tooth margin.

Thermocycling procedure

The samples were kept for 24 h in distilled water and placed in a thermocycler (Marconi, MA 470/3C, Piracicaba, SP, Brazil) at 5°C, 37°C, and 55°C in 500 cycles of 30 s in each bath, considering a transfer time of 10 s between baths. The samples were subjected to S. mutans biofilm activity as described below.

Induction of S. mutans biofilm formation

The blocks were covered with two layers of acid-resistant enamel (Colorama), except for half of the restored buccal surface (Fig. 1). No excess sealant had been observed at the preparation margin (optical microscope, 10x). Thus, half of the blocks, including enamel and sealant, were covered with acid-resistant varnish to create a control area (RI, EM, and CO) and an experimental area (RI-EX, EM-EX, and CO-EX), the latter of which was exposed to the biofilm.

In this phase, the samples were fixed in 45 wells (subdivided into 15 per group), in acrylic plates (TPP 24 ZellkulturTestplatte F, Trasadingen, Switzerland), and sterilized by ethylene oxide. The
mature biofilm was formed by a cariogenic bacterium, *Streptococcus mutans* ATCC 25175 (American type culture collection, Fiocruz, Rio de Janeiro, RJ, Brazil) that had been previously grown for 24 h in a brain heart infusion broth (BHI, Difco, Sparks, MD, USA) at 37°C under microaerophilic conditions. The bacterial inoculum was adjusted to the 0.5 McFarland standard. Subsequently, this bacterial suspension was diluted 1:100 and 10 µL was inoculated in each well with BHI supplemented with 2% sucrose. The plates were then incubated at 37°C under microaerophilic conditions for 48 h. The growth medium was renewed every 24 h.

**Knoop microhardness test**
The samples were positioned parallel to the coordinated table and adjusted to set the focus. Two rows of five indentations were performed with a Knoop indenter (HMV–G 21°, Shimada Microdurometer Corp., Japan) using a force of 0.05 HK (50 g or 490.3 mM) and an indentation time of 15 s (Fig. 1). The first indentation was performed at 25 µm from the interface and the other indentations kept a distance of 50 µm between them. The test was then performed. The same procedure was repeated for the control area after removing the acid-resistant varnish.

**Analysis of the microbial metabolic activity**
Microbial metabolic activity was measured by a colorimetric assay, which verified the enzymatic reduction reaction of the thiazolyl blue tetrazolium bromide (MTT, Sigma Aldrich, Sr. Louis, MO, USA) by *S. mutans* viable cells from the formed biofilm. The cylindrical samples (measuring 4 mm in diameter and 3 mm in thickness) were carefully prepared by dispensing the material in a standardized acrylic matrix. The insertion of the material and photopolymerization followed the manufacturers’ recommendation, as described in Table 1. These specimens (n=5) were transferred to cell culture plates (24 well plates) for the induction of *S. mutans* biofilm growth. *S. mutans* biofilm was induced according to the previously described methodology.

After 48 h of biofilm formation, the samples were carefully transferred to other sterile plates and 500 µL of MTT (1 mg/mL in PBS) was added to each well and the plates were then incubated at 37°C under microaerophilic conditions for 1 h in the absence of light. Subsequently, 500 µL of dimethyl sulfoxide (DMSO) was added to each well and the plates were gently mixed and incubated for another 20 min in the absence of light. The solutions were transferred to other plates and placed in a microplate reader at a wavelength of 540 nm. Higher absorbance indicates a higher metabolic activity of biofilm-forming cells.

**Statistical analysis**
The data were analyzed by GraphPad Prism, version 8.0 (GraphPad Software, Inc., USA). All data were initially verified using the Shapiro-Wilk test. Based on these preliminary analyses, the Knoop microhardness test data were subjected to the Kruskal-Wallis and Mann-Whitney tests. The control of Knoop microhardness after exposure to biofilm in the same groups was analyzed using Wilcoxon’s test.

All analyses were performed at a significance level of 0.05. Fig. 2 prepared by GraphPad Prism, version 8.0, presents the results for hardness and the coefficient of variation in the groups at different distances. The medians, minimum and maximum values, and coefficient of variation of the metabolic activity test results obtained for the *S. mutans* biofilm formed on the materials after 48 h are shown in Table 2.

**Results**

**Knoop microhardness test**
The average Knoop microhardness values and confidence interval for the marginal seal at different distances are shown in Fig. 2. Smaller values are observed for the enamel margins subjected to the biofilm when compared to the corresponding control. There was loss of hardness in all groups subjected to the cariogenic biofilm; however, this difference was not significant in the RI-EX group when compared to the RI control (paired Wilcoxon’s test; p>0.05). Regarding the extent of demineralization, a statistically significant difference was observed only at a distance of 25 m from the enamel/material margin in the areas exposed to *S. mutans* biofilm in all groups. At this distance, the RI-EX group showed the highest Knoop microhardness value compared to the CO-EX and EM-EX groups (Kruskal-Wallis; Mann-Whitney; p<0.05).

**Table 2.** Metabolic activity of *S. mutans* biofilm formed on the materials after 48 h

<table>
<thead>
<tr>
<th></th>
<th>CO</th>
<th>EM</th>
<th>RI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>0.061b</td>
<td>0.049a</td>
<td>0.055a</td>
</tr>
<tr>
<td>Minimum-maximum</td>
<td>0.059-0.065</td>
<td>0.040-0.052</td>
<td>0.053-0.061</td>
</tr>
<tr>
<td>Coefficient of variation</td>
<td>4.95%</td>
<td>13.29%</td>
<td>7.39%</td>
</tr>
</tbody>
</table>

Different letters indicate significant difference (p<0.05).
All materials showed accumulation of *S. mutans* biofilm after 48 h. The EM group showed a statistically significant difference when compared to the CO group; however, there was no statistically significant difference when compared to the RI group (Table 2).

**Discussion**

The present *in vitro* study evaluated the influence of two fluoride-releasing materials used in pit and fissure sealants for the prevention of enamel demineralization by *S. mutans* biofilm. Knoop microhardness and metabolic activity tests were used for the analysis. Several studies have used different methodologies to compare the application of ionomeric and resinous materials to pits and fissures in an attempt to prevent demineralization.[17,20-22] Therefore, in this study, a cavity preparation was performed to simulate the conditions of pits and fissures and the Knoop microhardness test was carried out to assess demineralization around the preparation margins after 48-hour biofilm formation. Knoop microhardness has been widely used for this purpose[18,23,24] and the *S. mutans* biofilm metabolic activity test, which can simulate oral conditions, has been used to replace cyclic pH.[22] Bovine teeth were used in this study. Human teeth are better at mimicking the effects of demineralization but obtaining the ideal number of teeth with a surface that is large enough to explore would pose a great challenge. Both histological and morphological similarities led to the choice of bovine teeth which, due to their large size and availability, allowed widely exploring the same enamel block, if necessary. Accordingly, bovine teeth are widely used in dental research.[25-27]

The antimicrobial activity and remineralization associated with the presence of fluoride-releasing sealants are widely acknowledged in laboratory and clinical studies.[4,18,28,29] Retention in and penetration into pits and fissures, risk of isolated carious lesions, and the inherent properties of the materials are some of the factors that can affect the effectiveness of a sealant.[19,20,21]

In the present study, fluoride-releasing materials interfered with the metabolic activity of the biofilm, confirming the antimicrobial activity of fluoride.[24] The model used in this study for prevention of demineralization might have contributed to the results found for the dental materials tested. Although both fluoride-releasing materials could prevent demineralization, the RI group showed better results as it interfered with the demineralization process at the tooth/material interface compared to the other material tested, and was therefore compatible with the performance of ionomeric materials described in other studies.[18,23] These results can be explained by the quick effect of fluoride-releasing ionomeric materials within the first hours, acting initially at the tooth/material interface and remaining constant at lower concentrations.[30] In addition, prevention of demineralization occurred after aging of the material. Aging by thermocycling, as used in this study, might have contributed to lower bacterial adhesion. According to Bürgers et al[29] aging can increase the material’s roughness but decrease free surface energy. The decrease in surface energy influences bacterial adhesion. Bürgers et al[29] found lower bacterial adhesion in several resinous and ionomeric materials, but they did not evaluate changes in enamel caused by the decrease in surface energy.

The results in this study corroborate those obtained by Amaral et al[18] in their *in situ* study, which showed greater remineralization efficiency in fissures treated with ionomeric sealant when compared to resinous materials without fluoride release. There were dentin carious lesions in 7.8% and 6.7% of teeth treated with glass ionomer cements and EWB, respectively.[31] Therefore, no statistical differences were observed in retention rates in any of the groups.[31] Note that, in the present study, enamel microhardness around materials was measured after a 48 h exposure to cariogenic challenge (*S. mutans* biofilm with culture medium supplemented with sucrose). The cariogenic effects tend to persist for a long time, as also demonstrated by the *in vivo* studies described above. EM has previously shown greater *S. mutans* inhibition compared to other fluoride-releasing materials and still maintained inhibitory
capacity for a long time.[32] In that same study, greater inhibitory capacity was obtained in sampled materials and not when associated with enamel.[32] Similar results were found in the present study. Thus, EM showed microbial inhibition when the material was tested immediately, but fluoride release could not be maintained after aging.

Regarding the accumulation of S. mutans biofilm on the surface of the materials, the CO group showed higher accumulation, whereas the EM group showed lower biofilm formation compared with the RI group, but there was no statistical difference between them. This may have probably occurred because of the larger fluoride release in the EM group. A large amount of fluoride was released in the first few hours, decreasing over time, possibly because of the closed polymer matrix formed after photopolymerization.[10]

On the other hand, glass ionomer cement maintained constant fluoride release for a long time.[10] Besides, EM also showed higher antimicrobial activity than other sealants with lower fluoride release.[10] This suggests that fluoride release may have been a relevant factor in the present study. In addition to the presence of fluoride, other factors can influence biofilm formation on the material's surface, such as surface roughness, chemical composition, and particle size,[33] but these factors were not tested in the present study.

Metabolic activity of S. mutans biofilm viable cells was quantitatively and indirectly measured on a spectrophotometer after the addition of MTT, which allows enzymatic reaction and changes the color of the medium. Even though this method does not quantify biofilm accumulation thoroughly (considering cell formation and extracellular matrix), it is widely used in several studies.[34-36]

This study has some limitations. The metabolic activity test assessed only S. mutans biofilm, which does not represent biofilm present in the oral cavity. Therefore, our results cannot be extended to clinical situations without some restrictions.[29] Furthermore, the Knoop microhardness test is a widely used indirect method for evaluating demineralization, but other methods can be used for the same purpose.[18,37-39] Evaluating the subsurface lesion formed could provide greater knowledge about the influence of these materials on the remineralization process or demineralization inhibition.

Interestingly, both fluoride-releasing materials were able to prevent demineralization even after the material was “aged” through the thermocycling process. Our study did not evaluate demineralization immediately after placement of the material, as in other studies.[17,39]

**Conclusion**

Our findings, albeit limited, show prevention occurred after the initial period of largest fluoride release. This underscores the importance of the aging process to interpret in vivo results.

In other words, the glass ionomer modified by resin has not been compared or evaluated in any other study regarding its ability to prevent caries, making comparisons between results impossible. Many studies have been carried out since the introduction of glass ionomers as pit and fissure sealants and their effectiveness has been questioned because of their low retention. In studies with resin-reinforced glass ionomer, better retention has been observed with fluoride release, but effectiveness is still unclear. The fluoride “refill” capacity from glass ionomer has been well known as a clinical advantage of the material, but this should be further investigated in future studies. Analyzing the results obtained for Knoop microhardness, metabolic activity tests demonstrated a greater ability to prevent demineralization in the enamel adjacent to the material in the RI group when compared to CO and EM groups. This prevention occurred close to the tooth/material interface at about 25 µm. Both fluoride-releasing dental materials (RI and EM) showed an inhibitory capacity against the metabolic activity of the cariogenic biofilm tested.

**Conflict of Interest:** None declared.

**References**


